

# Hepatoprotective effects of systemic ER activation

BulkRNAseq - Differential expression analysis

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```
# library import
library(tidyverse)
library(DESeq2)
library(edgeR)
```

## Load data

```
# removed outlier sample PPT_HFD_male_4 for differential expression (see PCA plot, fig. 1C)
# raw counts RNAseq
raw_counts <- read.table(
  file = 'data/bulkRNAseq_mmus_rawcounts.tsv',
  stringsAsFactors = FALSE,
  sep = '\t',
  header = TRUE) %>%
  dplyr::select(-PPT_HFD_male_4) %>%
  tibble::column_to_rownames('geneID') %>%
  as.matrix()

# design RNAseq
design_meta <- read.table(
  file = 'data/bulkRNAseq_mmus_design.tsv',
  stringsAsFactors = FALSE,
  sep = '\t',
  header = TRUE) %>%
  filter(sample != 'PPT_HFD_male_4')

# ensembl gene annotation (Mus musculus)
gene_ann <- read.table(
  file = 'data/ensembl_mmus_sep2019_annotation.tsv',
  stringsAsFactors = FALSE,
  sep = '\t',
  header = TRUE,
  fill = FALSE,
  quote = '')
```

# Differential expression

## Run DESeq2 pipeline

```
ds_data <- DESeqDataSetFromMatrix(countData = raw_counts,
                                  colData = design_meta,
                                  design = ~ 0 + condition)

ds_data <- estimateSizeFactors(ds_data)
ds_data <- DESeq(ds_data)

# filtering to minimum 15 counts in at least 8 samples
ds_data <- ds_data[rowSums(counts(ds_data, normalized=TRUE) >= 15) >= 8, ]

# annotations for gene background
gene_ids_bg <- rownames(counts(ds_data, normalized=TRUE))
gene_ann_bg <- gene_ann %>%
  filter(ensembl_gene_id %in% gene_ids_bg)

# comparisons
DESeq2_DEGs <- list(
  CDfVsCDm = results(ds_data, contrast = c('condition', 'CDf', 'CDm')),
  HFDfVsHFDm = results(ds_data, contrast = c('condition', 'HFDf', 'HFDm')),
  CDfVsHFDf = results(ds_data, contrast = c('condition', 'CDf', 'HFDf')),
  CDmVsHFDm = results(ds_data, contrast = c('condition', 'CDm', 'HFDm')),
  DPNVsHFDm = results(ds_data, contrast = c('condition', 'DPN', 'HFDm')),
  DIPVsHFDm = results(ds_data, contrast = c('condition', 'DIP', 'HFDm')),
  E2VsHFDm = results(ds_data, contrast = c('condition', 'E2', 'HFDm')),
  PPTVsHFDm = results(ds_data, contrast = c('condition', 'PPT', 'HFDm'))
)

# add annotation to DEG lists
DESeq2_DEGs <- lapply(DESeq2_DEGs, as.data.frame)
DESeq2_DEGs <- lapply(DESeq2_DEGs, rownames_to_column, var = 'ensembl_gene_id')
DESeq2_DEGs <- lapply(DESeq2_DEGs, inner_join, y = gene_ann, by = 'ensembl_gene_id')
```

## Run edgeR pipeline

```
groups <- design_meta$condition
dge <- DGEList(raw_counts, group = groups)
design <- model.matrix(~0 + groups)

# filter on CPM
dge <- dge[(rowSums(cpm(dge) > 1) >= 8), ]

y_dge <- calcNormFactors(dge, method = 'TMM')
y_dge <- estimateGLMCommonDisp(y_dge, design)
y_dge <- estimateGLMTrendedDisp(y_dge, design)
y_dge <- estimateGLMTagwiseDisp(y_dge, design)

fit_dge <- glmFit(y_dge, design)

# comparisons
edgeR_DEGs <- list(
```

```

CDfVsCDm = glmLRT(fit_dge, contrast = makeContrasts(groupsCDf-groupsCDm, levels = design)),
HFDfVsSHFDm = glmLRT(fit_dge, contrast = makeContrasts(groupsHFDf-groupsSHFDm, levels = design)),
CDfVsSHFDf = glmLRT(fit_dge, contrast = makeContrasts(groupsCDf-groupsSHFDf, levels = design)),
CDmVsSHFDm = glmLRT(fit_dge, contrast = makeContrasts(groupsCDm-groupsSHFDm, levels = design)),
DPNVsSHFDm = glmLRT(fit_dge, contrast = makeContrasts(groupsDPN-groupsSHFDm, levels = design)),
DIPVsSHFDm = glmLRT(fit_dge, contrast = makeContrasts(groupsDIP-groupsSHFDm, levels = design)),
E2VsSHFDm = glmLRT(fit_dge, contrast = makeContrasts(groupsE2-groupsSHFDm, levels = design)),
PPTVsSHFDm = glmLRT(fit_dge, contrast = makeContrasts(groupsPPT-groupsSHFDm, levels = design))
)

# calculate fdr and add annotation to DEG lists
edgeR_DEGs <- lapply(edgeR_DEGs, function(x) as.data.frame(x$table))
edgeR_DEGs <- lapply(edgeR_DEGs, rownames_to_column, var = 'ensembl_gene_id')
edgeR_DEGs <- lapply(edgeR_DEGs, function(x) mutate(x, padj = p.adjust(PValue, method = 'fdr')))
edgeR_DEGs <- lapply(edgeR_DEGs, inner_join, y = gene_ann, by = 'ensembl_gene_id')

```

## Common DEGs (DESeq2 & edgeR)

```

# filter DESeq2 results
DESeq2_DEGs_filt <- lapply(DESeq2_DEGs, function(x) filter(x, abs(log2FoldChange) > log2(1.75) & padj < 0.05))

# filter edgeR results
edgeR_DEGs_filt <- lapply(edgeR_DEGs, function(x) filter(x, abs(logFC) > log2(1.75) & padj < 0.05))

# intersect DEGs
common_DEGs_filt <- mapply(function(a, b) filter(a, ensembl_gene_id %in% b$ensembl_gene_id), DESeq2_DEGs_filt, edgeR_DEGs_filt)

```

## Export DEGs

```

DEGs <- list(unfilt = DESeq2_DEGs,
            filt = common_DEGs_filt)

saveRDS(DEGs, file = 'results/bulkRNAseq_mmus_DEGs.rds')

```

```
sessionInfo()
```

```

## R version 4.0.5 (2021-03-31)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows 10 x64 (build 19044)
##
## Matrix products: default
##
## locale:
## [1] LC_COLLATE=English_United States.1252
## [2] LC_CTYPE=English_United States.1252
## [3] LC_MONETARY=English_United States.1252
## [4] LC_NUMERIC=C
## [5] LC_TIME=English_United States.1252
##
## attached base packages:
## [1] parallel stats4 stats graphics grDevices utils datasets
## [8] methods base

```

```
##
## other attached packages:
## [1] edgeR_3.30.3          limma_3.44.3
## [3] DESeq2_1.28.1         SummarizedExperiment_1.18.2
## [5] DelayedArray_0.14.1   matrixStats_0.58.0
## [7] Biobase_2.48.0        GenomicRanges_1.40.0
## [9] GenomeInfoDb_1.24.2   IRanges_2.22.2
## [11] S4Vectors_0.26.1      BiocGenerics_0.34.0
## [13] forcats_0.5.1         stringr_1.4.0
## [15] dplyr_1.0.3           purrr_0.3.4
## [17] readr_1.4.0           tidyr_1.2.0
## [19] tibble_3.1.4          ggplot2_3.3.3
## [21] tidyverse_1.3.0
##
## loaded via a namespace (and not attached):
## [1] bitops_1.0-6          fs_1.5.0              lubridate_1.7.9.2
## [4] bit64_4.0.5           RColorBrewer_1.1-2    httr_1.4.2
## [7] tools_4.0.5           backports_1.2.1       utf8_1.1.4
## [10] R6_2.5.0              DBI_1.1.1             colorspace_2.0-0
## [13] withr_2.4.1           tidysselect_1.1.0     bit_4.0.4
## [16] compiler_4.0.5        cli_2.3.0             rvest_0.3.6
## [19] xml2_1.3.2            scales_1.1.1          genefilter_1.70.0
## [22] digest_0.6.27         rmarkdown_2.14        XVector_0.28.0
## [25] pkgconfig_2.0.3       htmltools_0.5.2       dbplyr_2.0.0
## [28] fastmap_1.1.0         rlang_0.4.10          readxl_1.3.1
## [31] rstudioapi_0.13       RSQLite_2.2.3         generics_0.1.2
## [34] jsonlite_1.7.2        BiocParallel_1.22.0   RCurl_1.98-1.2
## [37] magrittr_2.0.1        GenomeInfoDbData_1.2.3 Matrix_1.3-2
## [40] Rcpp_1.0.7            munsell_0.5.0         fansi_0.4.2
## [43] lifecycle_0.2.0       stringi_1.5.3         yaml_2.2.1
## [46] zlibbioc_1.34.0       grid_4.0.5            blob_1.2.1
## [49] crayon_1.4.0          lattice_0.20-41       splines_4.0.5
## [52] haven_2.3.1           annotate_1.66.0        hms_1.0.0
## [55] locfit_1.5-9.4        knitr_1.31            pillar_1.6.2
## [58] geneplotter_1.66.0    reprex_1.0.0          XML_3.99-0.5
## [61] glue_1.4.2            evaluate_0.14         modelr_0.1.8
## [64] vctrs_0.3.8           cellranger_1.1.0      gtable_0.3.0
## [67] assertthat_0.2.1      cachem_1.0.3          xfun_0.31
## [70] xtable_1.8-4          broom_0.7.4           survival_3.2-7
## [73] AnnotationDbi_1.50.3  memoise_2.0.0         ellipsis_0.3.2
```